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Extracellular alpha/beta-hydrolase from *Paenibacillus* species shares structural and functional homology to Tobacco Salicylic Acid Binding Protein 2

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Highlights:

- Identification of extracellular esterase from *Paenibacillus* species, PnbE.
- PnbE functions as an acetylsterase with optimum pH of 8.
- Crystal structure of PnbE shares structural homology to *N. tabacum* SABP2.
- Initial functional studies support PnbE functions as methyl salicylate esterase.

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Abstract

An alpha/ beta hydrolase annotated as a putative salicylate esterase within the genome of a species of *Paenibacillus* previously identified from differential and selective growth on Kraft lignin was structurally and functionally characterised. Feruloyl esterases are key to the degradation of lignin in several bacterial species and although this activity was investigated, no such activity was observed. The crystal structure of the *Paenibacillus* esterase, here denoted as PnbE, was determined at 1.32 Å resolution, showing high similarity to *Nicotiana tabacum* salicylic acid binding protein 2 from the protein database. Structural similarities between these two structures across the core domains and key catalytic residues were observed, with superposition of catalytic residues giving an RMSD of 0.5 Å across equivalent Ca atoms. Conversely, the cap domains of PnbE and *Nicotiana tabacum* SABP2 showed greater divergence with decreased flexibility in the PnbE cap structure. Activity of PnbE as a putative methyl salicylate esterase was supported with binding studies showing affinity for salicylic acid and functional studies showing methyl salicylate esterase activity. We hypothesise that this activity could enrich *Paenibacillus* sp. within the rhizosphere by increasing salicylic acid concentrations within the soil.

Keywords: alpha/beta-hydrolase; methyl salicylate esterase; *Paenibacillus*; rhizosphere; Salicylic Acid Binding Protein 2 (SABP2); salicylic acid

Database: structural data are available in the PDB under the accession number 6TJ2.

³ List of Abbreviations: PnbE, *Paenibacillus* esterase; SABP, salicylic acid binding protein; Nt, *Nicotiana tabacum*; LB, Luria broth; PMSF, phenylmethylsulfonyl fluoride; PEG 3350, polyethylene glycol 3350; RMSD, root mean square deviation.

1 Introduction

2 Lignin, an aromatic heteropolymer, makes up 15-30% of plant cell wall lignocellulose found
3 in plant biomass. The conversion of lignin into renewable chemicals as part of lignin-based
4 biorefinery is a current area of scientific interest (Bugg and Rahmanpour, 2015; Zakzeski et
5 al., 2010). Fungal degradation of lignin is well characterised throughout literature (Dashtban
6 et al., 2010; Hatakka, 1994; ten Have and Teunissen, 2001), however, fungal enzymes are
7 challenging to overexpress for industrial application. Bacterial breakdown of lignocellulose
8 from plant biomass requires a range of different extracellular enzymes including Dyp-type
9 peroxidases (Ahmad et al., 2011; Brown et al., 2012; Rahmanpour et al., 2016), multicopper
10 oxidases (Granja-Travez et al., 2018; Majumdar et al., 2014), and in one case an extracellular
11 manganese superoxide dismutase (Rashid et al., 2015). Multiple esterases have also been
12 shown to be important for either the hydrolysis of ferulate-arabinose linkages within
13 arabinoxylan (Millar et al., 2017; Topakas et al., 2007; Wong, 2006) or the hydrolysis of
14 xylan acetyl groups (Degrassi et al., 1998; Kosugi et al., 2002; Shao and Wiegel, 1995; Till et
15 al., 2013).

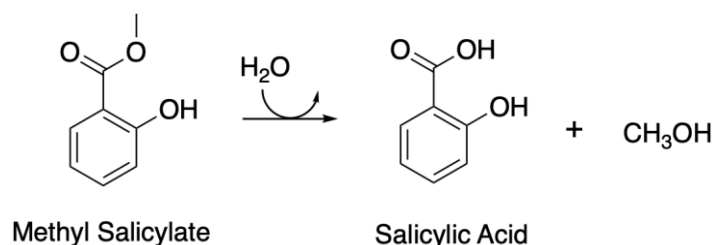
16 A wide variety of extracellular esterases have previously been implicated in the degradation
17 of lignin (Frago et al., 2016; Hong et al., 2012; Millar et al., 2017; Raymer et al., 1990).
18 Feruloyl esterases hydrolyse the ester linkages between ferulic acid and hemicellulose, with
19 this activity important as straw lignins can contain up to 5% ferulic acid (Buranov and
20 Mazza, 2008). Ferulic acid esterases have already been identified from *Aspergillus niger*
21 (Faulds and Williamson, 1994; Record et al., 2003), *Pseudomonas fluorescens* (Bartolome et
22 al., 1997) and *Streptomyces avermitilis* (Garcia et al., 1998).

23 Many bacterial species have already been shown to be involved in lignin degradation
24 including *Ochrobactrum* sp. (Granja-Travez et al., 2018), *Sphingobacterium* sp. (Rashid et
25 al., 2015), *Bacillus* sp. (Chandra et al., 2007; Degrassi et al., 1995), *Clostridium* sp. (Rogers

et al., 1992; Wang et al., 2013) and *Paenibacillus* sp. (Mathews et al., 2016; Song et al., 2014). A *Paenibacillus* sp. was isolated from an environmental soil sample by selective and differential growth on Kraft lignin, with genome sequencing indicating similarity to the characterised *Paenibacillus* sp. FSL H7-0737 (Granja-Travez et al., 2018). A multicopper oxidase from this species was previously investigated as a putative polyphenol oxidase enzyme (Granja-Travez et al., 2018). Further investigation into other extracellular enzymes from this species is therefore required to identify how it is utilising Kraft lignin.

Bioinformatic analysis of the genome of this *Paenibacillus* species identified an extracellular esterase (NCBI Reference Sequence: WP_076283022), here denoted as PnbE, is described as a putative salicylate esterase. Esterases represent a diverse group of hydrolases capable of catalysing the cleavage and formation of ester bonds. The majority of esterases belong to the carboxyl esterase gene family within the α/β -hydrolase fold protein superfamily (Hotelier et al., 2004; Montella et al., 2012). **α/β -hydrolase** fold domains are found across a range of functionally diverse enzymes, with a plethora of different substrates. They share a conserved β -sheet core of five to eight strands joined by α -helices in an $\alpha/\beta/\alpha$ sandwich format (Hotelier et al., 2004). PnbE shares 25% sequence homology with the methyl salicylate esterase (SABP2) from tobacco. Salicylic acid binding proteins (SABPs) were originally identified from protein extracts from tobacco leaves (Du and Klessig, 1997), with more recent studies using screening methods for identification (Manohar et al., 2014). Previously characterised salicylic acid binding proteins include carbonic anhydrases (SABP3 (Slaymaker et al., 2002)) and methyl salicylate esterases (SABP2 (Forouhar et al., 2005)). Methyl salicylate esterases catalyse the hydrolysis of the methyl group to form salicylic acid from methyl salicylate (Scheme 1). SABP2 has been implicated as important in both local and systemic resistance in plants (Kumar and Klessig, 2003; Tripathi et al., 2010). Enzymes catalysing release of

salicylic acid are of particular interest due to the prominence of salicylic acid in plant immunity (Durner et al., 1997) and root microbiome regulation (Lebeis et al., 2015).



Scheme 1. Conversion of methyl salicylate to salicylic acid.

In order to explore the function of this esterase in more detail, we have expressed this enzyme, and we report the functional and structural characterisation of the recombinant enzyme. We show that PnbE shows activity as an acetylcysteine esterase with a range of *p*-nitrophenol esters and is functional as a methyl salicylate esterase but shows no activity as a feruloyl esterase. We describe the structural similarities and differences of PnbE and NtSABP2. This is the first structural deposition of a bacterial esterase shown to have methyl salicylate esterase activity.

Materials and Methods

Cloning of PNB esterase. Genomic DNA from *Paenibacillus sp.* was extracted using Wizard® Genomic DNA Purification Kit (Promega). Forward and reverse primers were designed for a truncated esterase gene without the signal peptide to improve the expression. The forward primer (5' CACCGCC GCT AAC CAC AAA TCC TCT ACC AAA CAG3') and reverse primer (5' CTA ATC TCT ACC CGC CTT GAC CAT C 3') were designed for PCR amplification using Platinum Pfx-DNA polymerase (Invitrogen) as per manufacturer's instructions. The amplified gene was cloned using the Champion™ pET 151 Directional TOPO® Expression Kit (Invitrogen) into expression vector pET151 and transformed into *E. coli* TOP10 competent cells (Invitrogen). Purified plasmid (pET151::PnbE) was sequenced to confirm ligation and accuracy of the sequence.

Expression and Purification. Recombinant hexa-histidine tagged *Paenibacillus* esterase (PnbE) was expressed in *E. coli* BL21 cells (Invitrogen). To 2 L LB broth containing 100 µg/mL ampicillin a 1% inoculum of overnight starter culture of *E. coli* BL21 cells containing pET151::PnbE was added. The culture was then incubated at 37 °C, 180 rpm until an OD₆₀₀ of 0.6 was reached. Expression was induced by the addition of 0.5 mM IPTG followed by incubation at 15 °C overnight. Cells were harvested by centrifugation at 4000 ×g at 4 °C for 20 minutes. PnbE was purified using Nickel affinity chromatography. Harvested cells were resuspended in 20 mL lysis buffer (50 mM sodium phosphate pH 8.0, 300 mM NaCl, 1 mM PMSF). Cells were lysed with a constant system cell disrupter and debris pelleted by centrifugation at 10,000 ×g at 4 °C for 30 minutes. Cleared lysate was then loaded onto a Ni-NTA column (HisTrap HP, 1 mL) which had been pre-equilibrated in wash buffer (50 mM sodium phosphate pH 8, 300 mM NaCl, 20 mM Imidazole). The column was then washed with 100 mL of wash buffer before the protein was eluted at 0.5 ml/min with elution buffer

(50 mM sodium phosphate pH 8.0, 300 mM NaCl, 250 mM imidazole). PnbE was further purified using size exclusion chromatography with a Sephadex G-75 column (Sigma-Aldrich, St Louis, MO, USA). The column was equilibrated in 50 mM sodium phosphate pH 7.4, 100 mM NaCl at 0.75 mL/min.

PnbE Activity with Esters. PnbE activity assays with *p*-Nitrophenyl ester substrates (*p*-nitrophenyl benzoate, *p*-nitrophenyl palmitate, *p*-nitrophenyl phosphate, *p*-nitrophenyl acetate, *p*-nitrophenyl octanoate) were set up in duplicate in 50 mM sodium phosphate pH 8.0, 300 mM NaCl with 6 μ M PnbE and 2mM of the substrate. Reactions were monitored at 25 °C by a Cary 50 spectrophotometer by the appearance of *p*-nitrophenol at 405 nm (ϵ 18,000 M⁻¹ cm⁻¹) over time.

PnbE pH-rate Profile. The pH-rate profile for PnbE was determined at 25 °C using the assay described previously with *p*-nitrophenyl acetate (1 mM) as the substrate over a range of pH buffers. The buffers used contained 50 mM boric acid, 50 mM sodium acetate and 50 mM potassium phosphate and were then adjusted to the desired pH (every 0.5 unit between 3.5 and 10.5).

Analytical Size Exclusion Chromatography. A Superose 6 column (10 mm \times 300 mm, GE Healthcare) was used to estimate the apparent molecular mass of purified PnbE (~ 1 mg/mL). Protein samples were applied (1 mL/min) to a pre-equilibrated column in 50 mM sodium phosphate pH 7.4, 150 mM NaCl. Apoferritin (443 kDa), β -amylase (200 kDa), alcohol dehydrogenase (150 kDa), bovine serum albumin (66 kDa) and carbonic anhydrase (29 kDa) were used as protein standards for calibration of the column. Elution of the protein samples were monitored by absorbance at 280 nm.

Binding of PnbE to Salicylic Acid. Binding of salicylic acid to PnbE was determined using thermal shift assays. Reactions with 5 \times Sypro Orange, 10 μ M of PnbE and salicylic acid (at final concentrations of 0 to 4 mM) were made up in 10 mM NaH₂PO₄, 50 mM NaCl pH 8 in

a 96-well low profile white skirted plate (ThermoFisher). The plates were sealed with sealing films (EXCEL Scientific) and heated in an Agilent Technologies Stratagene MX300 5P Real Time PCR Detection System from 25 to 100 °C in increments of 0.1 °C. Fluorescence changes in the wells of the plate were monitored with the wavelengths for excitation and emission at 490 nm and 575 nm respectively. Binding curve was calculated using the single site ligand binding equation described by Vivoli *et al.* (Vivoli et al., 2014).

PnbE Salicylate Esterase Activity. Reaction contained 0.2 mg/mL PnbE and 10 mM methyl salicylate and were incubated at room temperature for 30 minutes. The reaction was stopped by incubating at 100 °C for 5 minutes. The reaction was then analysed by HPLC using Phenomenex Luna 5 µm C18 reverse phase column (100 Å, 50 mm, 4.6 mm) on a Hewlett–Packard Series 1100 analyser, at a flow rate of 0.5 ml/min, with monitoring at 210 nm. The gradient was as follows: 10 to 30% MeOH/H₂O over 10 min, 30 to 40% MeOH/H₂O from 10 to 20 min, 40 to 70% MeOH/H₂O from 20 to 30 min and 70 to 100% MeOH/H₂O from 30 to 40 min. Peaks corresponding to methyl salicylate or salicylic acid were identified by HPLC analysis of known standards (Fig. A.1).

Crystallization, data collection, and structure determination. Pure recombinant PnbE (21 mg/mL) in 50 mM sodium phosphate buffer pH 7.4, 150 mM NaCl was subjected to crystallisation screening using a Mosquito liquid handling robot (TTP Labtech). Protein (200 nL) was mixed with 200 nL of crystallisation solution from commercially available screens in MRC 96-well 2-drop crystallisation plates (Molecular Dimensions). Plates were sealed with sealing films (EXCEL Scientific) and incubated at 22 °C. Crystals appeared in a number of conditions between 1-2 weeks. Rod shaped crystals were grown in condition E1 of the PactPremier crystallisation screen (Molecular Dimensions) containing 0.2 M sodium fluoride, 20% PEG 3350. Crystals were removed from drops using a mounted Litholoop (Molecular Dimensions), cryoprotected in crystallisation solution containing 20% ethylene glycol, and

flash-frozen in liquid nitrogen. X-ray diffraction data to a resolution of 1.32 Å were collected at 100K at the beam line I03 at the Diamond Light Source, U.K. using a Pilatus 6M detector. All data were indexed, integrated and scaled using the XDS package (Kabsch, 2010). Further data handling was carried out using the CCP4 software package (Dodson et al., 1997). The structure was solved by molecular replacement using the automated pipeline by BALBES (Long et al., 2008), with the primary search model as PDB: 3DQZ. The structure was further refined by alternate cycles of manual refitting using Coot (Emsley and Cowtan, 2004) and Refmac (Murshudov et al., 1997). Water molecules were added to the atomic model automatically using ARP (Langer et al., 2008), at the positions of large positive peaks in the difference electron density, only at places where the resulting water molecule fell into an appropriate hydrogen bonding environment. Restrained isotropic temperature factor refinements were carried out for each individual atom with overall anisotropic scaling. Data collection and refinement statistics are given in Table 1.

Molecular Dynamics and Molecular Docking. The protein structure of PnbE (6TJ2) determined by x-ray crystallography was used for computational study. Missing N-terminal amino acids were input based on the known sequence of the protein using Accelrys discovery studio. After minimising and equilibrating the structure, 10 ns conventional molecular dynamics (cMD) simulations followed by 500 ns accelerated molecular dynamics (aMD) simulations were done using AMBER 16 package program (Case et al., 2005). MD analysis was used to extract the top 10 dominant conformation structures for running molecular docking. AutoDock SMINA (Koes et al., 2013) was used for blind docking of methyl salicylate and salicylic acid to determine the most probable binding site. GOLD (Jones et al., 1997, 1995) molecular docking was then used to determine the best-docked pose for each system, based on low binding energy and related score (Table A3.). Molecular dynamic

1 simulations, as described previously, for SABP2 (PDB:1Y7I) was also completed for
2 determination of flexibility and fluctuations for comparison with PnbE.

3 **Accession Number.** The co-ordinates and structure factors of the characterised *Paenibacillus*
4 sp. esterase has been deposited in the PDB with the accession code 6TJ2.

5

Results

Identification and Purification of *Paenibacillus* Esterase (PnbE). *Paenibacillus* sp. was isolated from a soil sample collected from Hampton Lucy, Warwickshire and differentially grown on Kraft lignin and minimal media (Granja-Travez et al., 2018; Rashid et al., 2017). The genome was sequenced and explored esterase enzymes, with preference for those exported from the cell. The gene for PnbE from *Paenibacillus* sp. was chosen for further exploration as a putative extracellular esterase. The sequence for PnbE is currently identified as an α/β -hydrolase by BLAST with the sequence ID of WP_076283022.

To improve overexpression of PnbE the sequence encoding the signal peptide was excluded by designing primers for PCR amplification of the gene without the signal peptide. Boundaries for the signal peptide were identified using SignalP-5.0 server (Nielsen et al., 1997) (Fig. A.2). The gene for PnbE was amplified from *Paenibacillus* 351 genomic DNA and cloned into a pET vector. The encoded protein is 253 amino acids in length (28.3 kDa) in the absence of the signal peptide. Expression in *E. coli* BL21 yielded high levels of PnbE (35 mg/L cell culture) which was then purified using metal affinity chromatography (Fig. A.3) and further purified by size exclusion chromatography.

Structure determination. The crystal structure of PnbE has been determined at 1.32 Å resolution (PDB accession 6TJ2). The asymmetric unit of the crystal contained three molecules with the space group P2₁2₁2₁ (Fig. 1a). The current atomic model contains residues 11-253 for the three chains within the asymmetric unit. Structural analysis shows a core domain with typical α/β -hydrolase fold and an α -helical cap domain (Fig. 1b).

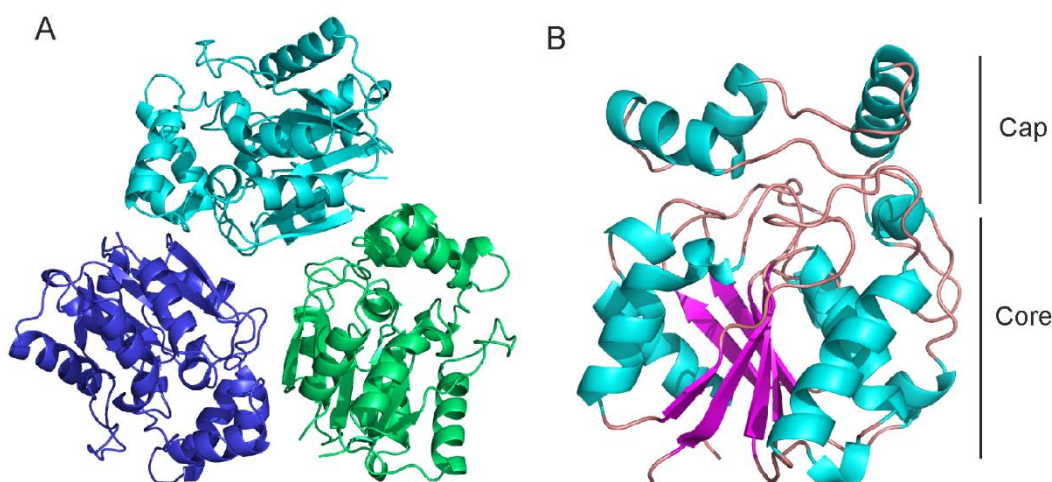


Figure 1. Overall structure of PnbE. A. Three chains present in the asymmetric unit. B. Subunit of PnbE, coloured by secondary structure (β -strands: magenta, α -helices: turquoise, loops: pale pink), with core and cap domains denoted.

Overall Structure. Three chains are observed within the asymmetric unit but PISA analysis (Krissinel and Henrick, 2007) suggested the absence of any meaningful interfaces between the subunits (Complex Formation Significance Scores of 0.000 between chains) (Table A.2). To further investigate this analytical size exclusion chromatography was used to confirm the multimeric state of the purified protein. PnbE formed one major peak with an apparent mass of ~150 kDa which suggests a hexamer (6×28.3 kDa, 170 kDa) (Fig. 2). Crystal structures for other α/β -hydrolase have also contained multiple chains within the asymmetric unit but have been shown to be active in the monomeric form (Forouhar et al., 2005; van Pouderoyen et al., 2001). We hypothesise that during purification and crystallisation the protein is at an unnaturally high concentration and therefore forms a multimeric structure of trimer or hexamer. Despite the presence of 10 hydrogen bonds and 3 salt bridges between non-crystallographic trimeric subunits within the ASU, PISA analysis (Krissinel and Henrick,

2007) concluded these did not have a role in complex formation but instead were a result of crystal packing. We therefore suggest that the active enzyme is monomeric.

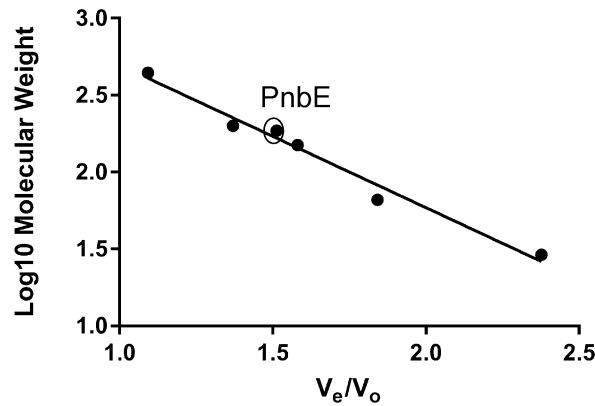


Figure 2. Multimeric state of PnbE established using analytical gel filtration. Position of PnbE with relation to standards (Apoferritin (443 kDa), β -amylase (200 kDa), alcohol dehydrogenase (150 kDa), bovine serum albumin (66 kDa) and carbonic anhydrase (29 kDa)).

Each subunit divides into two domains, a core domain (residues 11-115, 182-253) and a cap domain (residues 116-181) (Fig. 1b). The core domain contains six stranded β -sheet surrounded by eight α -helices, which fits the canonical structure of an α/β -hydrolases (Holmquist, 2000; Hotelier et al., 2004; Nardini and Dijkstra, 1999). The cap domain is made up of five α -helices.

Table 1. Crystallography data collection and refinement statistics.

Data collection^a	
Space group	P2 ₁ 2 ₁ 2 ₁
Cell dimensions [<i>a</i> , <i>b</i> , <i>c</i> (Å)]	47.86, 112.43, 126.85
Wavelength (Å)	0.91587
Resolution (Å)	57-1.32 (1.39-1.32)
Observations	1039504 (135229)
Unique reflections	159905 (10462)
<i>R</i> _{sym}	0.059 (0.739)
<i>I</i> /σ (<i>I</i>)	17.7 (2.0)
Completeness (%)	99.2 (94.7)
Redundancy	6.5 (6.1)
CC _{1/2}	0.999 (0.737)
Refinement	
<i>R</i> _{cryst}	0.161 (0.289)
Reflections used	153475 (10113)
<i>R</i> _{free}	0.183 (0.296)
Reflections used	6430 (449)
Non-hydrogen atoms	6444 (667 water molecules)
<i>B</i> -factor (Å ²)	
Protein	16.7
Water	27.3
Root-mean-squared deviation	
Bond lengths (Å)	0.015
Bond angles (°)	1.9
DPI coordinate error (Å)	0.048

^aValues in parentheses are for highest-resolution shell.

Structural Homology to SABP2. A sequence-based search within the protein database (PDB) showed highest similarity of PnbE to the *N. tabacum* salicylic acid binding protein (NtSABP2) (PDB: 1XKL) with 26% identity and 41% positives. This was supported by structural comparison using DALI (Holm, 2020) which showed highest similarity with a z-score of 26.1 for a hydroxynitrile lyase (PDB: 3DQZ) and a z-score of 25.3 for NtSABP2 (PDB: 1Y7I), with z-scores over three indicative of some structural similarity.

Superposition of NtSABP2 (PDB: 1XKL) and PnbE using SuperPose (Krissinel and Henrick, 2004) gave a local RMSD of 1.63 Å over Ca atoms within residues 3-48 (PnbE) and 11-56

(NtSABP2) (Fig. 4a). Superposition of PnbE with a hydroxynitrile lyase (HNL) from *Arabidopsis thaliana* (3DQZ) yielded a slightly higher local RMSD of 1.07 Å over Ca atoms within residues 3-46 (3DQZ) and 11-54 (PnbE) but a far worse global RMSD (of 9.38 Å vs 6.5 Å respectively). In addition, PnbE was found to contain a glycine (Gly20) and methionine (Met233) at corresponding positions of the threonine and lysine required for HNL activity (Padhi et al., 2010). Further analysis was therefore focussed on NtSABP2.

The core domains of PnbE and NtSABP2 share a high level of similarity with both containing six β-strands surrounded by eight α-helices (Fig. 3a). Despite the average number of residues in an α-helix is ~10 residues (Kumar and Bansal, 1998), of the eight α-helices in the core domain within PnbE three are 1-turn α-helices (average of 4 residues) whereas in NtSABP2 only two are 1 turn α-helices.

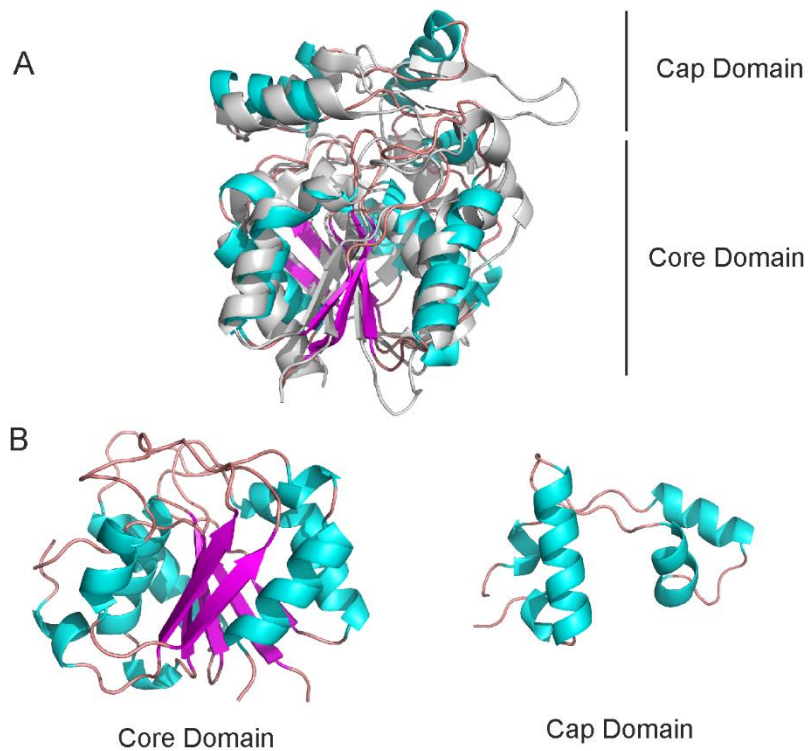
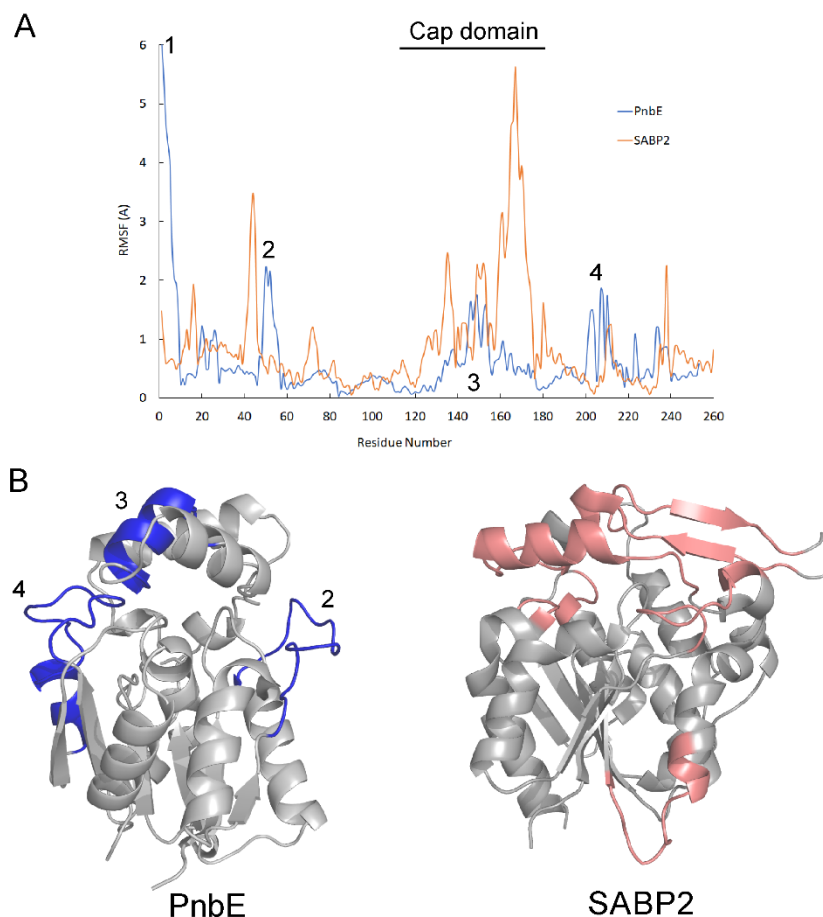


Figure 3. Structural similarities between PnbE and NtSABP2. A. Structural superposition of cartoon representations of PnbE and NtSABP2. B. Cartoon representation of core domain and cap domain for PnbE. PnbE coloured by secondary structure elements (β -strands: magenta, α -helices: turquoise, loops: pale pink) and NtSABP2 coloured grey.

The cap domains show greater divergence between NtSABP2 and PnbE. In NtSABP2 the cap domain is shown to contain three α -helices and two β -strands covering around 70 residues (Forouhar et al., 2005). Conversely, in PnbE there are five α -helices and no β -strands covering 67 residues (Fig. 3b). Within the cap domain there are two 1-turn α -helices at the hinge region of the cap domain as well as three that overlay well with those from NtSABP2 (Fig. 3a). Interestingly, RMSF plots indicate that the cap domain within PnbE (residues 116-181) shows reduced flexibility when compared to that within NtSABP2, with PnbE conversely showing greatest flexibility within the core domain (Fig. 4). Three flexible regions within the PnbE core domain were observed (Fig. 4b) with flexible region two containing the

1 residue Glu46, the oxyanion zone II residue which is essential for the stable formation of the
 2 oxyanion hole (Dimitriou et al., 2019). Furthermore, flexible region four contains the
 3 catalytic acid, Asp202, from the catalytic triad.



4 **Figure 4.** Predicted structural flexibility for PnbE and NtSABP2. A. Root Mean Square
 5 fluctuation (RMSF) plots for PnbE (6TJ2) and NtSABP2 (1XKL). B. Cartoon representation
 6 PnbE and NtSABP2 coloured grey, with flexible regions highlighted in blue and pink
 7 respectively. Flexible regions are numbered according the graph in 4a.

9
 10 CAVER analysis (Stourac et al., 2019) of both apo structures supports this with seven
 11 putative tunnels observed for NtSABP2 and only five putative tunnels observed for PnbE
 12 originating from the cavity around the active site. Comparison of the top throughput tunnel

for each structure shows an increase in length of the tunnel by $\sim 1.5\times$ and width of the tunnel bottleneck by $\sim 1.3\times$ comparatively between NtSABP2 and PnbE (Fig. A.4, Table A.3.).

Active Site. The majority of α/β -hydrolase contain a catalytic triad composed of a Nucleophile-His-Acid (Holmquist, 2000). In NtSABP2 these residues are Ser81, Asp210 and His238 (Forouhar et al., 2005) and from sequence and structural alignment comparable residues in PnbE are Ser85, Asp202 and His232. The catalytic triad residues are all situated in loops at the top of the core domain, with the putative active site enclosed by the cap domain (Fig. 5a).

PnbE appears to belong to group I (+++++) of the α/β -hydrolases classifications, as described by Dimitriou *et al.* in 2019 (Dimitriou et al., 2019). Other members of this group include SABP2, whereas AtHNL and HbHNL belonging to groups II and IV respectively. Comparison of the five key catalytic residues for PnbE, SABP2 and hydroxynitrile lyases are detailed in Table 2.

Table 2. Six key residues of the catalytic site surrounding the catalytic nucleophile.

	PnbE (PDB:6TJ2)	SABP2 (PDB:1XKL)	AtHNL (PDB:3DQZ)	HbHNL (PDB:3C6X)
X_{oxyII-2}	His19	His11	His11	His10
X_{oxyII-1}	Gly20	Gly12	Asn12	Thr11
W/Y/F	Trp28	Trp20	Trp20	Trp19
X_{ozII}	Glu46	Asp38	Glu38	Asp37
X_{nuc-1}	His84	His80	Phe80	Glu79
XIV	Asp109	Ala105	Asn105	Asn104

PnbE and NtSABP2 show strong conservation in the G-X-S-X-G motif, with both containing histidine at the nucleophile -1 position ($X_{\text{nuc}-1}$). Key residues involved in the formation and stabilisation of the oxyanion hole appear to be conserved between NtSABP2 and PnbE (Fig. 5b), including the oxyanion II -1 and -2 residues ($X_{\text{oxyII}-1}/X_{\text{oxyII}-2}$) and the tryptophan residue within the aromatic dipeptide located beneath the nucleophile and oxyanion hole (W/Y/F). The remaining aromatic residue in the aromatic dipeptide, however, differs between the two structures in both residue and position. The acidic residues at the oxyanion zone II position (X_{ozII}) also varies between the two structures, with glutamate and aspartate for PnbE and NtSABP2 respectively (Fig. 5b).

A/β-hydrolases can be further characterised by the positioning of the catalytic acid residue either at the end of strand β7 (Group A) or β6 (Group B) (Dimitriou et al., 2017). The coordinating position IV residue (XIV) located immediately after the strand β6 aids the coordination of the catalytic histidine loop above the plane of the catalytic acid zone (Dimitriou et al., 2017). Within PnbE the residue at position IV is Asp109, this is divergent from both NtSABP2 with the small non-polar residue Ala105 at this position and from AtHNL and HbHNL which contain Asn (Asn105 and Asn104 respectively). The presence of an acidic residue at this position is unusual and could implicate the formation of a structural catalytic tetrad, with both Asp109 and the catalytic acid Asp202 interacting with the catalytic histidine His85, rather than the classical catalytic triad previously observed for similar enzymes.

Superposition of the salicylic acid from NtSABP2 into the active site of PnbE shows close proximity to the catalytic triad residues, with around 3 Å distance from either His232 or Ser85 respective hydroxyl groups to SA comparable with distances observed for NtSABP2 (Fig. 5a). Positioning of the residues are very similar between the structures between the salicylic acid bound NtSABP2 (PDB:1Y7I) and the apo form of PnbE. Forouhar *et al.* previously showed that there was minimal difference observed between the ligand bound

1 (PDB:1Y7I) and apo form of NtSABP2 (PDB: 1XKL), with an RMSD of 0.45 Å between
2 equivalent Cα atoms (Forouhar et al., 2005). Superposition of the catalytic residues between
3 PnbE and NtSABP2 show a high level of similarity with an RMSD 0.50 Å between
4 equivalent Cα atoms.

5 To further explore the potential active site blind molecular docking was used to determine the
6 putative binding site, with ligands consistently occupying the pocket within the protein
7 containing the catalytic triad residues (Ser85, His232 and Asp202). Flexible molecular
8 docking of methyl salicylate (MSA) and salicylic acid (SA) using GOLD (Jones et al., 1997,
9 1995) in the SMINA-located binding site yielded good scores and energies of bindings for all
10 PnbE conformers tested (Table A.1). Figure 5c shows the 2D and 3D structure of best scored
11 docking of SA with a binding energy of -25.4 kcal/mol for SA and -24.43 kcal/mol for MSA.

12 Key residues found to bind SA include Asp109, Phe111, Phe212 and Ser85 (Fig. 5c) whereas
13 those for binding MSA include Ile205, Leu149, Phe111, Phe212 and Thr153 (Figure A.5).

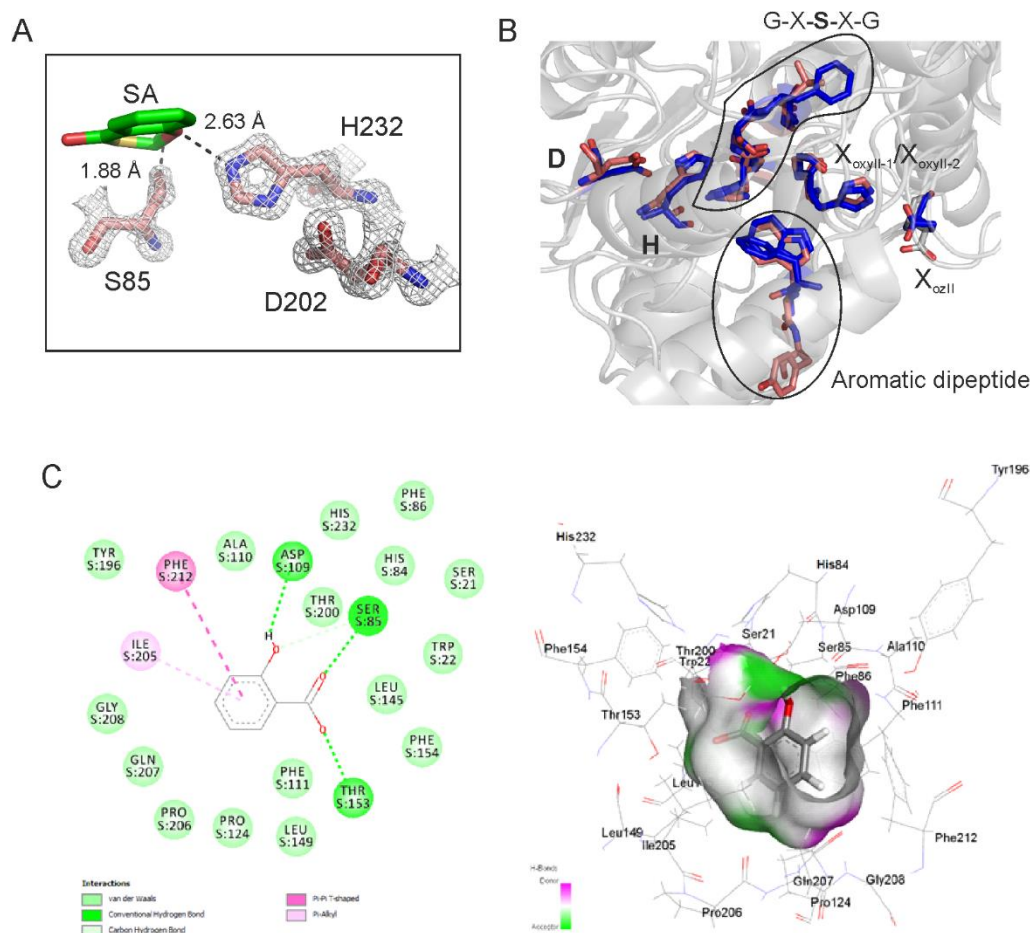


Figure 5. Putative Active Site for PnbE and similarity to NtSABP2. A. Superimposed SA in PnbE active site with catalytic triad residues displayed as pale pink sticks with electron density (σ level 2) shown and superposition of SA (displayed as green stick model) from NtSABP2 structure. B. superposition of key residues in PnbE active site (displayed as blue stick models) and corresponding residues in NtSABP2 (displayed as pale pink stick models), with catalytic triad residues are highlighted in bold. C. 2D and 3D structures of docked SA into the active site of PnbE. Key residues in ligand binding and the binding pocket are represented on the left and right respectively.

Esterase Activity. As PnbE is a previously uncharacterised esterase activity was originally tested with a range of different *p*-nitrophenol esters. PnbE was found to be active with *p*-nitrophenyl acetate (0.70 specific activity) and *p*-nitrophenyl octanoate (0.17 specific

activity) but showed no activity with *p*-nitrophenyl benzoate, *p*-nitrophenyl palmitate or *p*-nitrophenyl phosphate as a substrate (Fig. 6a). A pH-rate profile for PnbE with *p*-nitrophenyl acetate as a substrate showed the optimal pH for activity was pH 8.0 (Fig. 6b). Steady state kinetic parameters for PnbE with *p*-nitrophenyl acetate were calculated using a series of reaction time courses over a range of substrate concentrations (Fig. 6c). Analysis of reaction rates (V_0) vs. substrate concentration gave a calculated K_M of $627 \pm 53 \mu\text{M}$. These data suggest that the enzyme can function as an acetyl esterase with activity levels similar to those seen for previously characterised acetyl esterases (Millar et al., 2017; Watanabe et al., 2015).

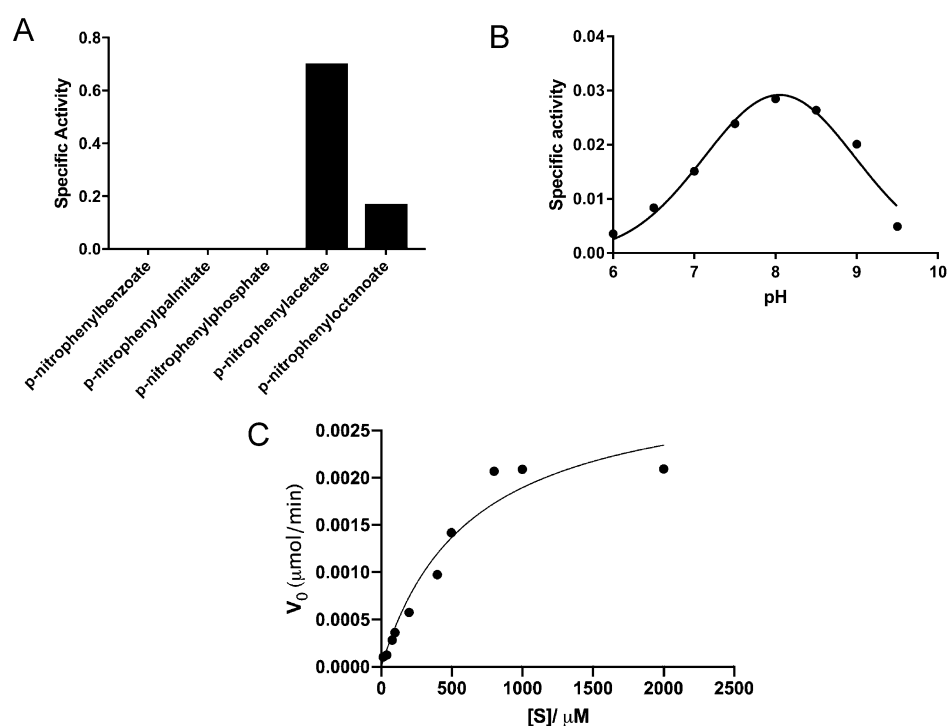


Figure 7. Esterase activity of PnbE. A. Specific esterase activity for PnbE in the presence of a range of putative substrates. B. pH profile for PnbE activity with *p*-nitrophenyl acetate. C. Michaelis-Menten plot for PnbE with 4-nitrophenyl acetate as a substrate.

Binding of salicylic acid (SA) to PnbE was determined using thermal shift assays (Huynh and Partch, 2015). The melting curves for PnbE shifted to the right with increasing concentration

of SA (Fig. 7a). The calculated melting temperature (T_m) for PnbE increased with the concentration of salicylic acid (Fig. 7b), confirming the ability of PnbE to bind SA.

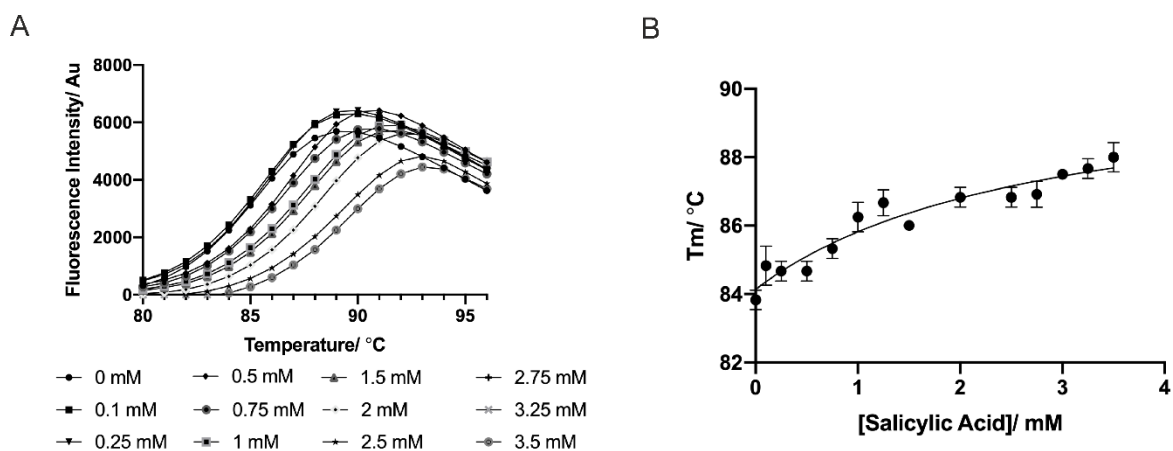


Figure 8. Binding affinity of PnbE for salicylic acid. A. Melting curves for PnbE with increasing concentrations of SA using thermal shift assays. B. SA binding curve for PnbE based on calculated melting temperatures (T_m) for PnbE over a range of SA concentrations.

With NtSABP2 detailed as a methyl salicylate esterase, the ability of PnbE to catalyse the hydrolysis of methyl salicylate to form salicylic acid was confirmed using HPLC analysis (Fig. 7b). Incubation of PnbE with methyl salicylate showed conversion to salicylic acid when compared to known standards, demonstrating that PnbE can function as a methyl salicylate esterase with a calculated k_{cat} of 0.40 s^{-1} . This can be compared to NtSABP2 which has a reported k_{cat} value of 0.45 s^{-1} (Forouhar et al., 2005).

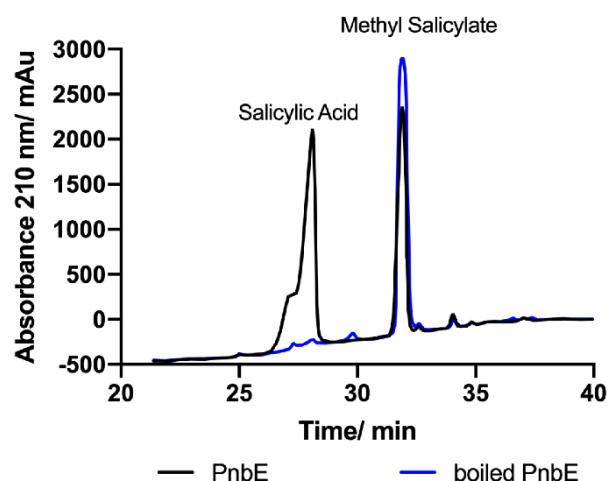


Figure 7. HPLC trace for the conversion of methyl salicylate to salicylic acid by 200 μ g PnbE (black line) in a 30 min assay (see Materials and Methods) compared to inactive (boiled) PnbE (blue line).

Discussion

Paenibacillus species can be found in both bulk soil (Uhlik et al., 2009) and the rhizosphere (Rosado et al., 1998; Timmusk et al., 2009) and have been identified as important in the degradation of lignin (Chandra et al., 2007). With limited information on the classes of extracellular enzymes involved in lignin degradation by *Paenibacillus* species we aimed to characterise the structure and function of PnbE, annotated as a salicylate esterase. We have verified activity as a methyl salicylate esterase, though PnbE shows no feruloyl esterase activity.

Sequence analysis of PnbE showed homology with *Nicotiana tabacum* Salicylic Acid Binding Protein 2 (NtSABP2), a methyl salicylate esterase. SABP2 enzymes have been shown to be important in local and systemic acquired resistance in plants (Kumar and Klessig, 2003; Tripathi et al., 2010). The phytohormone salicylic acid and derivatives are well known to be integral to plant immunity (Durner et al., 1997), with roles in cell response to infection as well as protection (Dieryckx et al., 2015; Kim et al., 2017).

1 Here we have described the determination of the structure of PnbE to 1.32 Å resolution which
2 can be classed as a group I α/β -hydrolase, according to the definitions by Dimitriou *et al.*
3 (Dimitriou *et al.*, 2019). Structural analysis using DALI (Holm, 2020) showed highest
4 similarity of the structure to that of a hydroxynitrile lyase (PDB:3DQZ) with a z-score of 26.1
5 and NtSABP2 (PDB:1Y7I) with a z-score of 25.3. Hydroxynitrile lyases (HNLs) and
6 esterases differ mechanistically and yet share key structural components such as the catalytic
7 triad (His-acid-nucleophile) and oxyanion hole (Rauwerdink and Kazlauskas, 2015). There
8 are however structural variances between the two, with HNLs containing a threonine which
9 blocks the entrance to the oxyanion hole via its side chain and a lysine within the active site
10 that can hydrogen bond with cyanide (Padhi *et al.*, 2010; Rauwerdink and Kazlauskas, 2015).
11 Mutations in these two residues has been shown to significantly reduce HNL activity in
12 *Hevea brasiliensis* HNL. Furthermore, esterase activity and HNL activity are antagonistic
13 and two mutations in NtSABP2 (Gly12Thr and Met239Lys) were sufficient to greatly reduce
14 esterase activity whilst introducing HNL activity (Padhi *et al.*, 2010). Structural analysis of
15 PnbE shows that it does not contain either of the residues linked to HNL activity, with
16 glycine and methionine at the corresponding positions respectively (Gly20 and Met233).
17 Other differences include that PnbE and HNL belong to different α/β hydrolase groups (I and
18 IV respectively). Therefore, despite marginally closer structural similarity to a hydroxynitrile
19 lyase, PnbE is unlikely to function as one. Conversely, PnbE shares several similarities with
20 NtSABP2 regarding key residues involved in the formation and stabilisation of the oxyanion
21 hole and catalytic activity, with catalytic triad residues superimposing with an RMSD of
22 0.498 Å between equivalent C α atoms. Regardless of the high levels of consensus between
23 the structures it is worth noting that sequence alignment showed that PnbE still aligns more
24 closely to other bacterial esterases than plant equivocal used for structural comparisons (Fig.
25 A.6).

1 Despite structural similarities between PnbE and NtSABP2 there are differences in their
2 substrate specificities, with lipase activity for *p*-nitrophenyl palmitate observed in NtSABP2
3 (Forouhar et al., 2005) but not in PnbE. Structural comparison using molecular dynamic
4 simulations showed different flexibility profiles, which could perhaps be unexpected as
5 proteins within the same fold families have been noted previously to have similar flexibility
6 profiles (Benson and Daggett, 2012). The reduced flexibility observed in the cap domain of
7 PnbE is corroborated by CAVER analysis (Stourac et al., 2019) showing fewer predicted
8 tunnels from the active site. Furthermore, the top throughput tunnel for PnbE is shorter and
9 narrower than that from NtSABP2. These data suggest that the active site in PnbE is less
10 accessible and this may explain the more stringent substrate specificity observed, particularly
11 for bulkier substrates such as *p*-nitrophenyl benzoate. The cap domains of α/β -hydrolases are
12 also known to contribute important residues to the active site (Rauwerdink and Kazlauskas,
13 2015), for PnbE there are two residues within the cap domain predicted to be involved in
14 binding SA or MSA, Leu149 and Thr153. Both of these residues are located within the only
15 area of increased flexibility within the cap domain of PnbE, suggesting that the conformation
16 of the cap domain is important in the binding of these compounds. Lipase activity is
17 dependent on seclusion of the active site from solvent (Bornscheuer, 2002) and five residues
18 from the cap domain were highlighted as shielding for SA in the active site of NtSABP2:
19 Asn123, Trp131, Phe136, Met149 and Leu181 (Forouhar et al., 2005). Comparative shielding
20 is not observed in PnbE (Fig. A.7.) and consequently this could account for the absence of
21 lipase activity in PnbE.

22 Confirmation of PnbE as a methyl salicylate esterase, originally based on sequence and
23 structural homology, using biochemical techniques showed binding of salicylic acid to PnbE
24 and conversion of methyl salicylate to salicylic acid. We therefore propose that PnbE serves
25 as a multi-functional extracellular esterase with acetyl esterase activity but more interestingly

1 catalytic activity for the conversion of methyl salicylate to salicylic acid with a calculated k_{cat}
2 of 0.4 s⁻¹. Methyl salicylate esterases have been described previously from Tobacco
3 (Forouhar et al., 2005), Arabidopsis (Vlot et al., 2008) and Poplar (Zhao et al., 2009) species
4 but there seems to be limited information for homologous activity within bacterial enzymes
5 and here we describe, to our knowledge, the first structure of a bacterial methyl salicylate
6 esterase.

7 The genera *Paenibacillus* is a member of the phylum Firmicutes, which have been previously
8 found within the rhizosphere (Mendes et al., 2013). Despite being isolated in both bulk soil
9 and rhizosphere, a recent study showed depleted levels of Firmicutes in bulk soil compared to
10 those in the rhizosphere for *Populus* species in Clatskanie soil (Veatch et al., 2019). Increased
11 salicylic acid concentration within the rhizosphere modulates the microbiome with an
12 observed increase the prevalence of Firmicutes and Actinobacteria species whilst decreased
13 prevalence of Bacteroidetes and Acidobacteria species (Lebeis et al., 2015). As a member of
14 the Firmicutes, *Paenibacillus* species, could be predicted to be at higher concentrations
15 within the rhizosphere and we propose that having an extracellular esterase capable of
16 releasing salicylic acid from methyl salicylate would therefore be beneficial. The means to
17 manipulate salicylic acid concentrations within the rhizosphere could allow for enrichment of
18 *Paenibacillus* species and other Firmicutes whilst reducing competition by depleting
19 Bacteroidetes and Acidobacteria species. We hypothesise this would likely be soil or plant
20 specific, as Veatch *et al.* conversely found a positive correlation between salicylic acid and the
21 acidobacterial Koribacteraceae and Solibacteria whilst describing no positive correlation with
22 Firmicutes for *Populus*-associated microbiomes (Veatch et al., 2019).

Conclusion

PnbE, an extracellular esterase from a *Paenibacillus* species differentially and selectively grown on Kraft lignin, shares structural similarities with the methyl salicylate esterase SABP2 from *Nicotiana tabacum*. This observation is compounded by functional studies which show binding of salicylic acid and catalytic turnover of methyl salicylate to salicylic acid. These data suggest a possible alternative function for PnbE to promote salicylic acid concentrations in the rhizosphere of plants which in turn could lead to enrichment of *Paenibacillus* species within the root microbiome.

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